

POZDYSHEV, V.A.; PENTIN, Yu.A.; TATEVSKIY, V.M.

Kinetics of the rotation isomerisation reaction. Vest. Mosk. un. Ser. mat., mekh., astron., fis. khim., 12 no.5:221-226 '57. (MIRA 11:9)

1. Laboratoriya molekuryarnoy spektroskopii Moskovskogo gosudarstvennogo universiteta.

(Isomerization)

APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

TATryckiv, VM.

USSR/Physical Chemistry - Kinetics, Combustion, Explosions, Topochemistry, Catalysis.

3-9

Abs Jour: Referat. Zmrnal Khimiya, No 3, 1958, 7221.

Author : V.A. Mal'tsev, A.D. Rusin, V.M. Tatevskiy.

Inst Title : To The Question of Spectroscopic Study of Combustion Reaction.

Orig Pub: Zh. fiz. khimii, 1957, 31, No 5, 1175-1176.

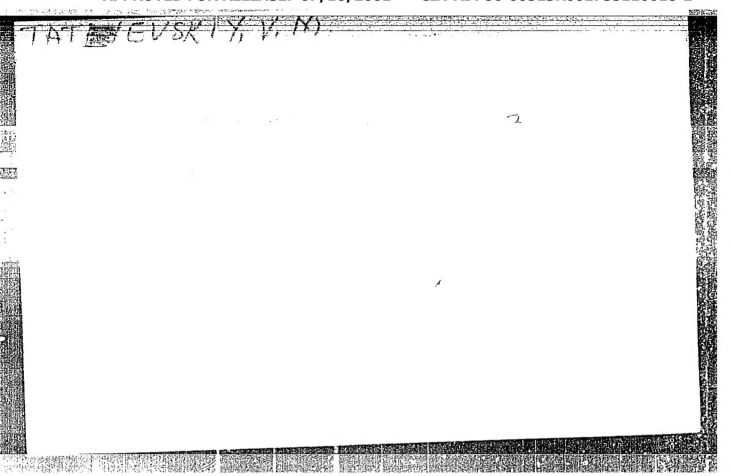
Abstract: The changes of the relative concentration of OH radicals in time are shown. These changes were recorded from the absorption spectrum at the explosion of H2 + 02 mixture on

a rapidly moving film.

Card : 1/1

-20-

APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

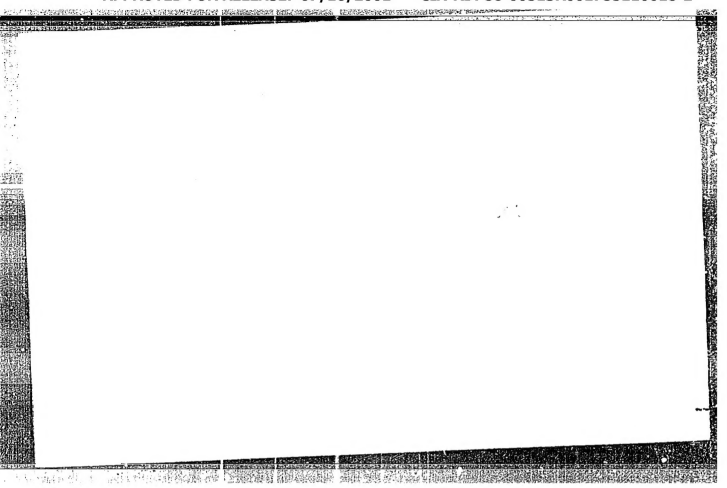


PENTIN, Tu.A.; TATEVSKIY, V.H.

Possibility of spectroscopic determination of the difference in energies of rotation isomers of complex molecules [with summary energies of rotation isomers of complex molecules [with summary in English]. Zhur.fiz.shiw.31 no.8:1830-1838 Ag '57'. (MIRA 10:12)

1. Moskovskiy gosudarstvennyy universitet im. M.Y. Lomonosova.

(Isomers—Spectra) (Molecules—Spectra)



NO A TOTAL PROPERTY OF THE PARTY OF THE PART PA - 2765 Chemical Structures and Physical and Chemical Properties of Hydrocarbons. Hew Laws and Calculation Methods. (Khimicheskoye stroyeniye ATTHOR: i fiziki-khimicheskiye svoystva uglevodorodov. Novyye zakonomernosti TITLE: Doklady Akademii Nauk SSSR, 1957, Vol 113, Nr 4, pp 836 - 838 Reviewed: 7 / 1957 PERIODICAL: (U.S.S.R.) Received: 6 / 1957 In the introduction to his paper the author cites some of his earlier publications. Equations are introduced which describe a number of physical and chemical properties of paraffins (alkanes). The same ABSTRACT: properties of paraffins of higher order are described by the equation $p_n^{(m)} = 2P_{12} - 3P_{22} + \Delta^{(m)} + nP_{22}$ shere n denotes the number of C-atoms in the paraffin, while the remaining values of the right part of the equation are constant, The coefficients of the straight line (5) can be determined if the values of P are known for two paraffins (alkanes) of the m-group and the values of P can be computed for all other paraffins of the same group (with different values of n) from this equation. In so far as the values P12 and P22 can be deterimend from the values of the corresponding physical and chemical properties of normal paraffins (alkanes) and $\triangle^{(m)}$ - from the value of the same physico-chemical Card 1/3

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Chemical Structures and Physical and Chemical Properties PA - 2765 of Hydrocarbons. New Laws and Calculation Methods.

property of one of the paraffins of the m-group, data on the properties of normal paraffins and the experimental value of the property P for one single paraffin of the same group can be used for the computation of the property under consideration of all paraffins of the group which was selected. Such methods of computing the physicochemical properties of paraffins of higher order as have hitherto been known are apparently extended and supplemented to a considerable degree by this equation. For all other classes of hydrocarbons with alkyle radicals the formula

$$P = C + \sum_{i < j=1}^{4} n_{ij}^{P}_{ij}$$
 (6)

was developed also previously. It can easily be seen that the hydrocarbons with a given core (C - const) can be divided into groups according to the character of the substituting alkyle radicals in such a minner that for every group (e.g. for the m-group) the equation can be transformed in two steps to give the final result $P_n^{(m)} = C_n^{(m)} + D_{22}^{(m)}$ (9)

$$P_n^{(m)} = C^{(m)} + \Delta^{(m)} + nP_{22}$$
 (9)

in which case all values except n are constant for the given group, and everything that has been said above holds also for hydrocarbons

Card 2/3

为19.77。19.71。19.71。19.11。19.11。19.12。19.12。19.12。19.12。19.12。19.12。19.12。19.12。19.12。19.12。19.12。19.12。19.12。1

Chemical Structures and Physical and Chemical Broperties PA - 2765 of Hydrocarbons. New Laws and Calculation Methods.

of other classes with alkyle radicals. Analyses confirm the predicitions of the theory. Furthermore, the author writes down equations for properties not subject to the equations (1) - (9) mentioned, as e.g. density and boiling temperature. For density the formula

 $d_{n}^{(n)} = \frac{a^{(n)} + b \cdot n}{g^{(n)} + h \cdot n}$

applies. With repsect to boiling temperature it is possible to show that it can be expressed in the case of the m-group of paraffins for a given pressure in first approximation by a fractured linear function. (5 citations from the author's publications)

ASSOCIATION: Moseow State University "M.V.LOMONOSOW"
PRESENTED BY: A.A.BALANDIN, Member of the Academy

SUBMITTED:

AVAILABLE: Library of Congress

Card 3/3

20-114-3-36/60

AUTHORS:

Pozdyshev. V. A., Pentin, Yu. A., Tatevskiy, V. M.

TITLE:

On the Kinetics of Reverse Isomerization Reaction (O kinetike

reaktsii povorotnoy izomerizatsii)

PERIODICAL:

Doklady Akademii Nauk SSSR,1957,Vol.114,Nr 3,pp 583-585(USSR)

ABSTRACT:

The potential barrier of the mutual transformation of the reverse isomers (transisomers or rotational isomers) of the molecules of the type of 1,2-dichloroethane, CXY, - CXY, can be represented by a formula found by Wolkenstein. Figure Nr 1 of the paper under review shows the shape of this function (curve Nr 1) for the transformation in the gaseous phase. In order to be able completely to determine the potential function, the three parameters V, V1, V2 would have to be known. This is necessary not only for the formation of a general concept on the nature of the phenomenon of retarded internal rotation and of the reverse isomerism of molecules, but also for the purpose of computing their thermodynamic functions. There exist no direct methods for the determination of the constants V , V of the retarding potential. In general, there was determined for the molecules of the type of 1,2-dichloro-

Card 1/3

20-111-3-36/60

On the Kinetics of Reverse Isomerization Reaction

ethane only a certain effective value of the retarding potential, for instance obtained from comparing the entropy value obtained on the basis of the third thermodynamic law and the value that had been computed under assumption of the free internal rotation or under simplified assumptions on the shape of the barrier ($V_2 = 0$, $V_1 = V$). It is possible to determine accurately, by means of experiments, one of the above parameters (V2). The authors of the paper under review attempt to demonstrate that in some cases it is also possible to determine the second parameter (V_4) by experiments. The value of 7188 can be measured with relatively high accuracy during investigation of the process of isomerization in the solid phase. Knowledge of the values V_4 and V_2 obviously also makes it possible to determine the third parameter of the potential curve V, if the value of the effective barrier is obtained from comparing the experimental entropy value and the value computed statistically under assumption of the free internal rotation. Thus the method of investigating the reaction kinetics of reverse isomerization as presented for the first time in the paper under review opens new and interesting prospects for research in the field of the phenomena

Card 2/3

	PRASE I BOCK EXPLOITATION SOW/1700		Are po spektroskopii, 1956. (Materials of the 10th All-Union Vol 2: Atomic Spectroscopy) 1958. 958 p. (Sattes Its: 3,000 eoples printed.	Mademiya nenk 250%, Komissiya po	0.3. Landsborg, Academician, (Resp. Mi.); (1) Destor of Physical and Mathematical Sciences; (2) Destor of Physical and Mathematical Sciences; (2) Destor of Physical and Mathematical Sciences; (2) Candidate of Technical Sciences; (2) Landsdate of Technical Sciences; (3) Landsdate of Technical Sciences; (4) Landsdate of Technical Sciences; (4) Landsdate of Technical Sciences; (4) Landsdate of Technical Sciences; (5) Landsdate of Technical Sciences; (6) Landsdate of Technical Sciences; (6) Landsdate of Technical Sciences; (7) Landsdate of Landsdate	#7	d technical studies All-Union Confer- exarted out by and include ourses. The	uction, physicochemical methods for controlling uction, physica and technology of gas discharge, pectroscopy, abnormal disperion in metal vegors, and the combustion theory, spectrum estalysts of oreal photocychic methods for quantitative spectrum estale and alloys, spectral description of the fact of an example of the entire of metals and alloys, spectral description of the entire of metals we are all the entire of metals and allows.	ark spectrographic eralysis, n in the parameter of calibration es of metals, spectrum analysis in n metallurgy, and principles and nalysis.	on Conference (Cont.) 807/1700	Studying the Photometrie 195	n. M. Ye. Miryusevich, and V.H. s in the Design of the Didition cas for the Turpose of Resolving 195 thius Resonance Line		Ogurtaova. Radiation From 199	A.V. Bedoupseov, and A.Ye. Movik, Effect is Admixture on Low-pressure Nertury 201	Eordrasheva, Concave Eirror Lisorption in Light Sources 204
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CIA-RDP86-00513R001755110018-2" APPROVED FOR RELEASE: 07/16/2001

TATEVSKIY, V. M. (MGU, Moscow)

V. M. Tatevskiy, Yu. A. Pentin, Ye. G. Treshchova, and Kh. Kesler, "Rotational Isomerism and the Energy of the Formation of Hydrocarbons."

report presented at the Symposium on Concepts of Conformation in Organic Chemistry which took place in Moscow at the IOKh AN SSSR (Institute of Organic Chemistry, AS USSR) from September 30 to October 2, 1958.

Izvestiya Akademii nauk SSR, Otdeleniye khimicheskikh nauk, 1959, No. 3, 561-564.

SOV, 156 -58-2-1/48 · AUTHORS: Akishin, P. A., Naumov, V. A. Tatevskiy, V. M Electron-Diffraction-Investigations of the Molecular Struc-TITLE: ture of Gallium-Halides (Elektronograficheskoye issledovaniye stro/eniya molekul galogenidov galliya) PERIODICAL: Nauchnyye doklady vysshey shkoly, Khimiya i khimicheskaya tekhnologiya,1958, Nr 2, pp 205-209 (USSR) ABSTRACT: The task of the present paper is the determination of the configuration and of the geometrical parameters of the molecules of the gallium-fluoride, chloride, and bromide by the diffraction-method with fast electrons, with a vapor-jet of the substance to be investigated. Because no data have been found so far in this field (except Ref 1) the authors planned to apply a thoroughly worked-out experimental method as well as a deciphering method. For this reason a new type of electron-diffraction camera was employed (Ref 2) and the ampulla was filled in a drying room. The elaboration of a vapor-electron-diffraction-pattern was carried out visually and photometrically (Ref 3). The process of decoding was carried out Card 1/3 by means of: a) construction of curves of the radial distri-

Electron-Diffraction-Investigations of the Molecular Structure of Gallium-

bution (Ref 4) and b) the method of successive approximation (Ref 5). These investigations disclosed a new insight into the molecular structure of gallium-halides. The electrondiffraction-method proved the presence of monomer-molecules in vapor. Moreover the configuration was determined as well as the geometrical parameter of the CaF,-molecule. The dimerisation of molecules was proved in vaporous gallium-chlorides and gallium-bromides, and more reliable data were obtained from their structure and their geometrical parameters (see table 2) There exist good reasons to assume analogy between structure of the crystalline gallium-halides and aluminium-halides. Data on the configuration of gas molecules of Ga2Cl6 and Ga2Br6 agree with data of solid aluminium, gallium, and indium halides (Ref 9) as well as solid trimethyl-aluminium. There are 3 figures, 2 tables, and 9 references, 2 of which are Soviet.

ASSOCIATION:

Card 2/3

Kafedra fizicheskoy khimii Moskovskogo gosudarstvennogo universiteta im. M. V. Lomonosova (Chair of Physical Chemistry of the State University imeni M. V. Lomonosov, Moscow)

• ,	Electron-Diffraction-Investigations of the Molecular Structure of (
	SUBMITTED:	October 21, 1957								
	Card 3/3									

SOV/120-58-4-24/30

AUTHORS: Rusin, A. D. and Tatevskiy, V. M.

TITLE: A Time-Scan Spectrograph (Spektrograf s razvertkoy vo vremeni)

PERIODICAL: Pribory i tekhnika eksperimenta, 1958, Nr 4, pp 101-102 (USSR)

ABSTRACT: A medium-quartz spectrograph is fitted with a rotating cylinder cassette, electrically interlocked with the shutter; the minimum rate of motion of the spectrum across the film is 500 mm/sec, the maximum 8000 mm/sec; time marks are inserted by chopping the incoming beam. The spectrograph is of aperture f/22 and is used in studies on H₂/O₂ explosions; the

pressure wave in the vessel and the spectra are correlated in time by using an auxiliary photocell coupled to the chopper to mark on the pressure record. The system can also be used

Card 1/2

SOV/120-53-4-24/30

• A Time-Scan Spectrograph

contains 2 figures, no tables and 7 references, 2 of which ASSOCIATION: Khimicheskiy fakul'tet MGU (Chemistry Dept., MGU)

SUBMITTED: July 17, 1957.

MAL'ITSEV, A.A.; KORYAZHKIN, V.A.; MISYUMEVICH, M.Te.; TATEVSKIY, V.M.

Some modifications in the design of the recording system of the DIS-4 spectrophotometer to detect the isotope shift in the resonance line of lithium. Fix.sbor. no.4:195-197 '56.

(MIRA 12:5)

1. Khimicheskiy fakul'tet Moskovskogo ordena Lenina gosudar-stvennogo universiteta ineni M.V. Lomonosova.

(Lithium-Spectra) (Spectrophotometer)

APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

5(4)

AUTHOR:

Tatevskiy. V. M.

SC7/55-58-4-26/31

TITLE:

New Regularities in the Physical-Chemical Properties of Hydrocar bons. IV. Linear Depondences Between Different Physical-Chemical Properties of Alkenes of Single Homologous Groups (Novyye zakonomernosti v fizike-khimichaskikh svoystvakh uglavodorodov. IV. Lissynyya zavisimosti mashin razlichnyri flaiko-khimicheskimi svoystvami alkanov otdel'nykh gomologicheskikh grupp)

PERIODICAL: Vestnik Moskovskogo universiteta, Seriye anworkytki, wekhoulini, astronomii, fiziki, khimii, 1958, Nr 4, pp 205-203 (USSE)

ABSTRACT:

The author / Ref 1,2,3 7, V.A. Kireyav / Ref 5 7, and M.Kh. Karapetyants / Ref 6,7 7 investigated theoretically the equation

 $P_n^{(m)} = A_{PQ}^{(m)} Q_n^{(m)} + B_{PQ}^{(m)}, A_{PQ}^{(m)} = const, B_{PQ}^{(m)} = const.$

which expresses the linear connection between two different properties P and Q in the given (m-th) homology group of alkanes.

Card 1/2

APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

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Hew Pegalarithes in the Physical-Gharical Properties of Hydrocarbons. IV. Linear Dependences Between Different Physical-Chamical Properties of Alkanes of Single

In the present paper the formula (1) is compared with experimental results; for some cases the constants A and P are given. The author discusses the applicabilities of (1). There are I figures, 1 table, and 9 Soviet references.

ASSCCIATION: Kafeira fizioheskoy khizii (Chair of Physical Chemistry) SUBMITTED: July 15, 1967

Card 2/2

IMLLUSKI AUTHORS: Kot; ov, Yu.I., Tyulin, V.I. TITE: an! Raman Spectrum of Gaseous CO2. (Spoktr kombinataiornogo rasseyaniya gazoobraznoy (0,.) PERIODICAL: Optika i Spektroskopiya, 1958, Vol.IV, Nr.2, pp.271-272 ABSTRACT: Using apparatus described earlier (Nef.3) the authors obtained rotational spectrum of CO2 at 1.5-2 atm pressure on a grating spectrometer with linear dispersion of 6.7 A/nm in the second order. The spectral width of the spectrometer slit was 0.45 cm-1. The figure on p.271 shows a photograph of the spectrum obtained. spectrum was used for calibration. The results are given in the table on p.272. For a linear CO2 molecule displacements of rotational lines relative to the exciting $|\Delta v| = (4B-6D)(J+3/2) - 3D(J+3/2)^2$ where B and D are rotational constants, and J is the motational quantum number. From the results obtained the constant D could not be determined, but one our say Card 1/2

Raman Spectrum of Gaseous CO2.

51-4-2-25/20

that it is less than 2×10^{-5} cm⁻¹. Weglecting D the author finds $B_0 = 0.3295 \pm 0.0004 \text{ cg}^{-1}$. value of Bo agroes well with the value found from infrared spectra (Ref.2, p.422). Values of the vibrational frequencies for fully-symmetric vibration vibrational frequencies for fully-symmetric vibration were also obtained. They were found to be 1205.7 and 1388.2 cm⁻¹; these values agree with the values given in Refs.4 and 5. There are 1 figure, 1 table and 5 references of which 1 is Soviet, 3 American and 1 a translation of American work into American.

ASSOCIATION: Moscow State University. (Moskovskiy Gos. universitet)

1. Raman spectrum-Carbon dioxide 2. Carbon dioxide-Spectrographic

Card 2/2

SOV/51-5-5-5/23

AU THORS:

Tatevskiy, V.M., Tunitskiy, L.W. and Novikov, M.M.

TITLE:

Vibrational Constants and Dissociation Energy of the BeF Molecule. (Kolebutel'nyye postcyannyye i energiya dissotsiatsii molekuly BeF)

PERIODICAL: Optika i Spektroskopiya, 1958, Vol 5, Nr 5, pp 520-529 (USSR)

ABS TRACT:

The paper reports a new investigation of the vibrational structure of the A2IT--K2T bands of BeF. The BeF spectrum was excited using gas-discharge tubes. Since the gas temperature is comparatively low in such tubes the rotational structure of the bands was weaker than in an arc and this reduced overlapping of bands and made it easier to observe new band edges. Two types of tubes were used in this work: one working (Fig 2). Both tubes were filled with helium under 2-3 mm Hg pressure. BeF molecules were introduced into the discharge tubes by placing some discharges was U-chaped (1, in Fig 1) and had a quartz window (7, in Fig 1). The space around the electrodes (5, in Fig 1) was joined to the tube supplied from a 630 W transformer at 13 kV. The pulse-discharge source

Card 1/3

Vibrational Constants and Dissociation Energy of the BeF Molecule

SOV/51-5-5-5/23

consisted of a straight tabe (1, in Fig 2) with a fluorite wind me (2, in Fig 2). Its electrodes (4, in Fig 2) were supplied every second (for 0.02 sec) with a 600-700 V, 9 Å rules. The spectra were recorded using a DFS-3 spectrograph of 2 Å/mm dispersion and 144000 resolving power. An iron spectrum was used as a wavelength standard. Hoasurements were made with 0.5 cm-1 precision. The spectrum is shown in Fig 3. The measured band edges are given in Table 1. 48 new edges of Q1-branches and 54 new edges of R2 and R1 branches were recorded. Table 2 gives the rotational constants ω_0 , $\omega_0 x_0$ and $\omega_0 y_0$ of BeF taken Table 3 gives the values of v_{max}^{*} , v_{max}^{*} , v_{max}^{*} , v_{max}^{*} , v_{max}^{*} , and v_{max}^{*} and v_{max}^{*} allowing anharmonicity $v_0 y_0$ of the BeF molecule. Table 4 gives the recommended, host reliable values of the spectroscopic constants of the BeF molecule

Card 2/3

SOV/51-5-5-5/23

Vibrational Constants and Dissociation Energy of the Bel Molecule

This table gives the dissociation energies of the $X^2\Sigma$ and $A^2\Pi$ states as 8 ± 0.5 and 3.9 ± 0.5 eV respectively. The author thanks L.V. Gurvich for his advice. There are 4 figures, 4 tables and 7 references.

SUBLITTED: December 6, 1957

1. Beryllium fluoride--Spectrographic analysis 2. Molecules--Spectra Card 3/3

3. Molecules--Energy 4. Gas dishcarges--Applications

SOV/51-5-5-9/23

Treshchova, Ye.G., Tatevskiy, V.M., Skwarchenko, V.R. and Levina, R.Ya. AUTHORS:

Raman Scattering Spectra of Hydrocarbons of Various Types. (Spektry TITLE: koabinatsionnogo rasseyaniya uglevodorodov razlichnykh klassov). V. Raman Scattering Spectra of Certain Bi- and Diene Hydrocarbons (V. spektry kombinatsionnogo rasseymiya nekotorykh

bi- i tritsiklicheskikh diyenovykh uglevodorodov).

PERIODICAL: Optika i Spektroskopiya, 1958, Vol 5, Nr 5, pp 553-560 (USSR)

The Ramar spectra of the following bi-cyclic diene hydrocarbons with ABSTRACT: isolated and conjugated bonds were obtained: 2,2'- and 1,1'-dicyclohexenyl, 2,2'- and 1,1'-dicyclopentenyl, and dicyclopentadiene. 2,2'-dicyclohexenyl and 2,2'-dicyclopentenyl were synthesized by the action of Mg on 3-chlorcyclohexene-l and 3-chlorcyclopentene-l, respectively. 1,1'-dicyclohexenyl and 1,1'-dicyclopentanyl were obtained by de-hydratton of the corresponding pinacols (pinacones). Dicyclopentadiene of technical grade was purified by double washing with a concentrated alkaline solution, by washing with water, drying and vacuum distillation.

The boiling point, vapour pressure, refractive index and density at Card 1/2 20°C, etc., are given for all the five hydrocarbons in Table 1. The

Raman Scattering Spectra of Hydrocarbons of Various Types. V. Raman Scattering Spectra of Certain Bi-Space and Tri-Cyclic Diene Hydrocarbons

Raman spectra of 2,21-dicyclopentenyl and of dicyclopentadiene were obtained using a glass "Shtaynkhel'" spectrograph, as described in Refs 1, 2. The Raman spectra of 2,2'-dicyclohexenyl (Fig 1 curve v), 1,1'-dicyclohexenyl (Fig 1 curve a) and of 1,1'-dicyclopentary! (Fig 1 curve b) were obtained using a triple-prism glass spectrograph ISP-51 with photoslectric recording. Table 2 gives the values of the Raman fraquencies and intensities of all the five hydrocarbons. Reproducibility of the results was satisfactory and the differences between individual measurements of the strong lines did not exceed 1 3% (see Fig 2). The results obtained are discussed and interpreted in detail. As part of the discussion the authors quote the C==C frequencies and intensities of various dienes with isolated and conjugated double bonds (Table 3). There are 2 figures, 3 tables and 12 references, 11 of which are Soviet and 1 German.

SUBMITTED: Docember 31, 1957

Card 2/2

1. Hydrocarbons--Spectra 2. Roman spectroscopy 3. Spectrophotometers --Performance

AUTHORS: Kuzyakov, Yu.Ya. and Tatevskiy, V.M.

SOV/51-5-6-10/19

TITLE:

New Bands of the CF Molecule (Novyye polosy molekuly CF)

PERIODICAL: Optika i Spektroskopiya, 1958, Vol 5, Nr 6, p 699 (USSR)

ABSTRACT:

Andrews and Barrow (Ref 1) found only 4 bands belonging to the $A^2Z-X^2\Pi$ transition of the CF molecule: (0,0), (0,1), (1,0), (1,1). Other bands of this system could not be observed because of strong overlapping by complex bands of the CF2 molecule (Refs 1, 2). Under more stable conditions of an electric discharge through a stream of CF4 vapour, the present authors were able to decrease considerably the CF₂ band intensities and to measure 9 more bands of the $\mathbb{A}^2\mathbf{I}-\mathbf{X}^2\Pi$ transition. Assuming that the edges of the new bands are due to the same branches as in the case of the 4 bards reported earlier, the authors calculated the positions of the zero lines of the 9 bands: (0.2), (0.3), (0.4), (0.5), (0.6), (1.4), (1.5), (1.6), (1.7), observed by the author for the first time and listed in col. 1 of a table on p 599. This table gives the zero lines found experimentally (col. 2) and calculated (col. 3) from the vibrational constants given in Ref 1. The good agreement between the experimental and calculated values confirms the correctness of the vibrational analysis. More precise

Card 1/2

New Bands of the CF Molecule

SOV/51-5-6-10/19

values of the zero lines and the vibrational constants will be published later, when the full analysis of the rotations is conclets. Measurements of the fine structure of the (0.2) and (0.3) tanks yield the following values of the rotational constants: $B_2^{\prime\prime}=1.340$ cm⁻¹ and $B_3^{\prime\prime}=1.341$ cm⁻¹. The absence of the transitions with $v^{\prime}=2$ both in the experiments reported in the present paper and those reported by other workers suggests pro-dissociation between vibrational levels $v^{\prime}=1$ and $v^{\prime}=2$ of the $A^{\prime\prime}\Sigma$ state. In this case the upper limit of the dissociation energy of the ground state of the CF molecule should not exceed 5.8 eV. This is a complete translation except for the table. References: (1) K.B. Andrews, R.F. Barrow, Proc. Phys. Soc. London, A64, 481, 1951. (2) P. Venkateswarlu. Phys. Rev., 77, 676, 1950.

SUBMITTED: April 12, 1958

Card 2/2

TATEVSKIY, V.H.

New aspects of the physicochemical properties of hydrocarbons. Part 4: Linear dependence of the various physicochemical properties of alkanes of different homologous series. Vegt. Mosk.un. Ser. mat., uekh., astron., fiz., khim. 13 no. 4: 205-208 | 58. (MIRA 12:4)

1. Kafedra fizicheskoy khimii Hoskovskogo universiteta. (Paraffins)

AUTHORS: Tatevskiy, V. H., Benderskiy, V. A.

sov/79-28-7-4/64

TITLE:

New Rules Governing the Physicochemical Properties of Hydrocarbons. II (Novyye zakonomernosti v fiziko-khimicheskikh svoystvakh uglevodorodov.II)

PERIODICAL:

Zhumnal obshchey khimii, 1958, Vol. 28, Nr 7, pp. 1733 - 1737 (USSR)

ABSTRACT:

In the present paper the authors show that the theory suggested by the authors (Refs 1-3) that all alkanes could be divided into groups according to the principle of the same structure of the ramified fragments of the hydrocarbon chains is actually proved experimentally by various physicochemical properties of table. According to this table for each groups are given in the physicochemical quantities typical of the group must depend the given group (Schemes 1 and 2, where P₂₂ denotes the constant coefficient, and Fig. 4). The table of the group denotes the constant

coefficient, and Fig 1). In the abbreviated formula (2) P denotes the physicochemical constant, m a group, n the number of carbon atoms, $a^{(m)}$ the constant for the given group and β that

Card 1/3

New Rules Governing the Physicochemical Properties of Hydrocarbons. 11.

SOV/79-28-7-4/64

for all groups. In figures 1-4 it is shown to which degree the linear/rules(1) and (2) in every alkane group for various physicochemical properties of the alkanes are satisfied, All rules to be seen from the table and the figures can easily be used for the calculation of the physicochemical properties of the hydrocarbons of the given group not to be investigated, as far as the corresponding experimental data for some of the hydrocarbons of the same group are given, which is also of practical importance. There are 6 figures, 1 table, and 5 refer-

ASSOCIATION: Moskovskiy gosudarstvennyy universitet (Moscow State University)

June 1, 1957

Card 2/3

AUTHOR: Tatevskiy, V. M. SOV/79-28-11-7/55 TITLE: New Rules Governing the Physico-Chemical Properties of Hydrocarbons (Novyye zakonomernosti v fiziko-khimicheskikh III. The Heats of Crystallization of n.-Alkanes (III. Teploty kristallizatsii n.-alkanov) PERIODICAL: Zhurnal obshchey khimii, 1958, Vol 28, Nr 11, pp 2935-2939 ABSTRACT: Earlier, V. M. Tatevskiy (Refs 1,2) showed that such properties of hydrocarbons as molecular volume and molecular fraction, heat of formation from atoms and elements, heat of combustion, heat of evaporation, logarithm of vapor pressure and others in every alkane group with a characteristic ramification of the carbon chain depend on the number n of the carbon atoms in the alkane compound. This paper deals with the problem of similar rules governing the crystallization heat of alkanes. Due to the insufficient data existing, and as only the n.-alkanes are an exception of this rule, the author had to restrict to the latter. In determining the Card 1/3 rules governing the heats of crystallization, difficulties

arise that are connected with the occurrence of the solid phase, which is not the case if the properties of the gaseous phase and the liquid phase can be determined. It is important that in the crystallization various crystalline modifications can occur. Simple rules between the heat of crystallization and the atomic number of carbon in the molecule can exist only if the heats of crystallization of the alkanes are in relation to one and the same modification. This fact is important in the establishment of rules governing the heats of crystallization of n.-alkanes, as their crystalline modification is different in dependence on the even or oid atomic number, as well as on other properties. Thus, the dependence of the heats of crystallization of the n.-alkanes on the number of carbon atoms in the molecule was investigated. It was shown that the heats of crystallization of the low members of the n.-alkane series linearly depend on the atomic number, but that these dependencies of the cdd and even numbers n of the carbon atoms do not agree as the corresponding hydrocarbons are crystallized in different modifications.

Card 2/3

New Rules Governing the Physico-Chemical Properties SOV/79-28-11-7/55 of Hydrocarbons.

III. The Heats of Crystallization of n.-Alkanes

The figures and tables enclosed illustrate the results of these investigations. There are 5 figures, 2 tables, and

ASSOCIATION:

Moskovskiy gosudarstvennyy universitet (Moscow State University)

SUBMITTED:

July 7, 1957

Card 3/3

TATEVERLY, V. M.

75-1-20/32

AUTHORS:

Dombrovskaya, U., Pentin, Yu. A., Dombrovskiy, Ya., Satevskiy, V. M., Kochetkov, H. K.

TITLE:

The Investigation of the Tautomerism of the Alkyl-f-Aminovinyl Ketones According to Infrared Absorption Spectra (Issledovanine tautomerii alkil-faminovinilketonov po infrakrasnym spuktram pogloshcheniya)

PERIODICAL:

Zhurnul Fizicheskoy Khimii, 1998, Vol. 32, Nr 1, pp. 139-140 (USSR)

ABSTRACT:

The present work is directly connected with earlier works (reference 1 and 2). The investigations in these works referred to the clarification of the question of the existence of enamino-inide--tautomerism. The present work is also devoted to the same question. The spectra obtained experimentally are discussed here and compared with each other. 1) The range 2000 - 3500 cm⁻¹, of the valence-X-H-oscillations. In the spectrum of the solid anyl-p-aninovinyl ketone, i.s. in the co-form of the montance absorption bands with 3140 and 3222 cm⁻¹ corresponding to the oscillations of the OH group, are present. It is shown that the co-form is a more

Card 1/3

or less pure imide-cnol form.
2) The range 1450 - 1700 cm-1 of the double bonds. All data obtained here speak in favour of the fact that the oc-form corresponds

APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

The Investigation of the Tautomerian of the Alk/1-f-Aminovinyl Ketones According to Infrared Absorption Spectra

to the imide-enol form, and that the transformation product of the ∞ -form and the mixture of the ∞- and p-form represents a mixture of the imide-enol- and of the enumine-hetche-form. 3) The range 700 - 1450 cm⁻¹. In the spectrum of the mixture of the &-1 and f-forms of the aethyl-f-aminovinyl ketone 1250 - and 1002 cm ban's are present, which were not observed in the spectrum of the &-form and are characteristic for the enamine-ketone-form. Generally, an interpretation of the bands of this range is very difficult. 4) The range 400 - 700 cm . In the spectrum of the mixture of ∞ - and β -forms of homologues of the alkyl- β -granovinyl hotoms side bands with an absorption sentre ~ 650 cm are prekotones wide bands with an absorption sentre ~ 650 ca sent. These bands become eguentially more intensive in the spectra of the &-form. It is assumed that those ones correspon! to the deformation escillations of the hydroxyl group of the imide-enol--form. It may be assumed that the conclusions drawn with respect to the other ranges also apply for this range. Summing up it is proved on the base of the infrared absorption spectra (in the range of 400-3500 ca⁻¹) of the methyl- β -aminovinyl hetone, of its homelogues and of some model-sempounds in different physical states that the ally1-\$-mainsviny1 hetones exist in tautomeric forms; viz. as an enumina-ketone-form (A) and as an

Card 2/3

The Investigation of the Tautomerism of the Albyl-E-Aminovinyl Ketones According to Infrared Absorption Spectra

imide-enol-form (B). There are 1 figure, and 15 references, 8 of which are Slavic.

ASSOCIATION: Mosecu State University ineni M. V. Lomonosov

(Moskevskiy gosudarstvenny universitet in. M. V. Lomonocova)

SUBMITTED: October 24, 1955

AVAILABLE: Library of Congress

Card 3/3

AUTHORS:

Ziborov, V. A., Pentin, Yu. A., Tatevskiy, V. M.

76-32-3-33/43

TITLE

Letters to the Editor (Pis'ms v redaktsiyu).
Rules Governing the Mean Magnetic Susceptibility of
Paraffin Hydrocarbons and Edathols (Zakonomernosti
v srednikh magnitnykh vospriimchivostyakh parafinovýkh
uglevodorodov i spirtov)

PERIODICAL:

Zhurmal Fisicheskoy Khimii, 1958, Vol. 32, Nr 3, Pp. 707-709 (USER)

ABSTRACT:

In commercian with work performed on types and subtypes of hydrocarbon bonds and with corresponding analysis of their physicochemical properties a formula is given by which the phenomena manticmed in the title can be determined. The most acceptance of the determinations performed with a series of substances are given in a table. They show good agreement with the experimentally obtained values. The given series of calculation is more suitable than Pascal's scheme (and the modifications), and moreover does not possess any additive

Card 1/2

Letters to the Editor Rules Governing the Mean Magnetic Susceptibility of Paraffin

ocustants. The symliad theoretical expressions of the stability of the physiconhanical quantities (with the nature of a capacity faction) can, according to the results in comparison with the experimental data, be well used. In this manuar the importance of the three controls of organic compounts can be emplained and producted on a much broaden basis that it was possesse before. There are 2 tables and 8 Seviation preview. The state of the s

ASSOCIATION:

Mckovakiy gradicestrancy universitat in. M.V. Lancerva (Mose of Shade University inent M. V. Interpetal)

STANFOURD:

60tob € 17, 1956

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1-11 17 621

AUTHOR: Tatevskiy, V. M. 76-32-5-37/47 TITLE: Linear Relationship Between Various Physico-Chemical Properties of Hydrocarbons (Lineynyye zakonomernosti mezhdu razlichnymi fiziko-khimicheskimi svoystvami uglevodorodov) PERIODICAL: Zhurnal fizicheskoy khimii, 1958, Vol. 32, Nr 5, pp.1168-1170 ABSTRACT: Already in the papers by M. Kh. Karapet'yants (Refs 1 - 3) a linear relation between two properties of certain groups of related compounds was found. The present work deals with the theme mentioned in the title and starts from the earlier developed theoretical representations. It was already proved that the hydrocarbons can be divided intogroups in which the ramification of the chain parts and the cycles are the same, with a number of properties depending linearly on the number of carbon atoms n in the groups. Starting from the case that in various hydrocarbon groups two different physico-chemical properties are placed next to each other and that they are measured at various temperatures, pressures etc., it is found Card 1/2 after a number of considerations and mathematical deductions

Linear Relationship Between Various Physico-Chemical Properties of Hydro-

that in the case of a change of temperature a comparison of the property values can not be carried out in the given way. There are 8 references, 8 of which are Soviet.

ASSOCIATION: Moskovskiy gosudarstvennyy universitet im. M. V. Lomonosova (Moscow State University imeni M. V. Lomonosov)

April 19, 1957 1. Hydrocarbons--Physical properties 2. Hydrocarbons---Chemical properties

Card 2/2

SUBMITTED:

AUTHOR:

Tatevskiy, V. M.

SOV/76-32-6-6/46

TITLE:

The Rules Governing the Structure and the Physico-Chemical Properties of Alkanes (Zakonomernosti v stroyenii i fiziko-

-khimicheskikh svoystvakh alkanov)

PERIODICAL:

Zhurnal fizicheskoy khimii, 1958, Vol. 32, Nr 6,

pp. 1226 - 1230 (USSR)

AND DESCRIPTION OF THE PERSON OF THE PERSON

ABSTRACT:

With an increase of the number of carbon atoms in hydrocarbons the number of possible isomers rises rapidly. For this reason A.P.Meshcheryakov proposed not to compute single isomers in investigations of alkanes, but families with the same number of carbon atoms and comparable values of the fundamental physical and chemical properties. In this connection two important problems emerge: 1) The determination of the families, the hydrocarbons of each class decompose. 2) The determination of the number of isomers pertaining to each family. In order to solve the first problem, two stages of consideration are adopted for alkanes. On the one hand the alkanes with a given number of carbon atoms are divided into groups, which possess the same number of primary, secondary, tertiary and quaternary C-atoms. On the other hand the alkanes with the possible C4-C4 bindings

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The Rules Governing the Structure and the Physico-Chemical Properties of Alkanes

SOV/76-32-6-6/46

of the bindings C_i - C_j of each sub-type is determined. The equations pertinent to the first stage are deduced, as well as those pertinent to the second stage. In the latter the problem of the relation of the carbon atom group of each group with the possible number of C-C bindings of the various sub-types, primary-primary, primary-secondary, primary-tertiary etc. is taken into consideration, thus permitting to determine the number of C-C bindings. The results obtained furnish a further contribution towards a solution of the problems of a bulk computation of the physical and chemical properties of higher alkanes. There are 9 references, 6 of which are Soviet.

ASSOCIATION: M

Moskovskiy gosudarstvennyyuniversitet im. M.V. Lomonosova

(Moscow State University imeni M.V.Lomonosov)

SUBMITTED:

October 17, 1956

Card 2/3

The Rules Governing the Structure and the Physico-Chemical Properties of Alkanes

SOY/76-32-6-6/46

1. Methanes--Physical properties 2. Methanes--Chemical properties 3. Methanes --Structural analysis 4. Methanes--Test results 5. Mathematics

Card 3/3

TATEVSKIY, V.M.

..JTHORS:

Akishin, P. A., Vilkov, L. V., Tatevskiy, V. M. 20-1-33/58

. ITLE:

Electron Diffraction Study of the Chloroprene Molecule (Elektronograficheskoye issledovaniye stroyeniya molekuly

khloroprena).

FERIODICAL:

Doklady AN SSSR 1958, Vol. 118, Nr 1, pp. 117-120 (USSR)

ABSTRACT:

The task of the present work is the determination of the spacial configuration and the geometric parameter of the chloroprene molecule by means of the method of the diffraction of quick electrons with a vapour jet of the substance to be investigated. The apparatus for the taking of electronograms was already described in a preliminary work (ref. 1). With long waves of electrons of from 0,0520 to 0,0540 % 7 series of electronograms were obtained. With these electrons 8 maxima and 7 minima were measured by means of visual evaluation of their intensity. From the experimental data obtained and given in a table the curve of radial distribution was constructed. The calculation made with a variation of the values of the intensity of the extremes proved the reliability of the curve of radial distribution. The attachment of the peaks of the curve of radial distribution to interatom distances in the chloroprene molecule arranged by the authors is

Card 1/3

Electron Diffraction Study of the Chloroprene Molecule.

20-1-33/58

mentioned here. Then the theoretic curves of intensity of various models (the structure parameters of which are mentioned in a table) are calculated according to the method of successive approximation. Not with all models the theoretic intensity curves coincide with the experimental curve of scattered electrons. This non-coincidence exists e.g. for the plain transisomer, the plain zis-isomer as well as for the rotated isomer with a rotation of the vinyl-groups by 90° around the C2 - C3 group. The best coincidence is obtained for the plain model of the chloroprene molecule with transposition of double compounds and $C_1C_2C_3$ — and $C_2C_3C_4$ — angles differing by 5° , as well as for the non-plain model of the molecule with a rotation of the vinyl-groups around the only C2-C3-compound by 32° (rotation from the trans-position) and with equal C1C2C3- and C2C3C4- angles. The main parameters of the two latter models coincided completely with the interatom distances obtained from the curve of radial distribution. The two most probable structures resulting from the electronographic investigation of the structure of the chloroprene molecule are given with their numeric parameters. The double carbon compounds in the chloroprene molecule have a transfiguration or a similar configuration, which coincides with

Card 2/3

Electron Diffraction Study of the Chloroprene Molecule.

20-1-33/58

the earlier obtained infrared- and ultraviolet spectra of

chloroprene. There are 3 figures, 2 tables, and 11 references,

3 of which are Slavic.

Moscow State University imeni M. V. Lomonosov (Moskovskiy ASSOCIATION:

gosudarstvennyy universitet imeni M. V. Lomonosova).

January 3, 1957, by N. N. Semenov, Academician. PRESENTED:

December 29, 1956 SUBMITTED:

Library of Congress AVAILABLE:

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Card 3/3

CIA-RDP86-00513R001755110018-2" APPROVED FOR RELEASE: 07/16/2001

24(7) SOV/156-59-2-1/48 AUTHORS: Akishin, P. A., Naumov, V. A., Tatevskiy, V. M. TITLE: The Electronographical Investigation of the Structure of the Molecules of the Neodymium Halogen Compounds (Elektronograficheskoye issledovaniye stroyeniya molekul galogenidov neodima) Nauchnyye doklady vysshey shkoly. Khimiya i khimicheskaya PERIODICAL: tekhnologiya, 1959, Nr 2, pp 229-232 (USSR) ABSTRACT: In a previous investigation it was found (Ref 1) that the molecules of LaX, (X = halogen) have a plane configuration with the lanthanum atom in the center of an equilateral triangle. Because of the similar structure of the outer electron shells a similar configuration was to be expected in the case of neodymium. The measurements carried out by means of an electronograph of the Chemical Department of the MGU (Moskovskiy gosudarstvennyy universitet - Moscow State University) confirm this. The electronograms were read according to the method of radial distribution and according to the method of successive approximation. Figure 1 shows the curves of radial distribution. The two peaks of the curve are interpreted as r(Nd - X) and r(X - X). They were in agreement with the expected plane Card 1/2 configuration. The RMS oscillation amplitudes were computed

SOV/156-59-2-1/48 The Electronographical Investigation of the Structure of the Molecules of the Neodymium Halogen Compounds

> and the theoretical curves of the scattering intensity were plotted (Fig 2) which are in good agreement with the experimentally found curves. Table 2 shows the RMS oscillation amplitudes for NdF_3 , NdCl_3 , NdBr_3 and NdJ_3 and the geometrical parameter. The experimentally found symmetrical configuration of LaX, (Ref 1), YX, (Ref 7) and now also NdX, confirm the quantum chemical assumptions (Ref 8). In the series chlorine bromine - iodine, neodymium - halogen follow approximately a linear law, whereas the distance neodymium - fluorine is considerably reduced, as it is the case with a number of fluorine compounds. There are 2 figures, 2 tables, and 10 references, 5 of which are Soviet.

PRESENTED BY: Kafedra fizicheskoy khimii Moskovskogo gosudarstvennogo universiteta im, M. V. Lomonosova (Chair of Physical Chemistry, Moscow State University imeni M. V. Lomonosov)

SUBMITTED:

October 13, 1958

Card 2/2

24(7) SOV/156-59-2-2/48 AUTHORS: Kuzyakov, Yu. Ya., Tatevskiy, V. M.

TITLE: On the Spectrum of the CC1-Molecule (O spektre molekuly CC1)

PERIODICAL: Nauchnyye doklady vysshey shkoly. Khimiya i khimicheskaya

tekhnologiya, 1959, Nr 2, pp 233-236 (USSR)

ABSTRACT: The oscillation analysis of the absorption bands of the electric discharge in carbon tetrachloride as suggested

by P. Venkatesvarlu (Ref 2) is rejected and a new oscillation analysis is given. Table 1 shows the measured spectral lines and the frequencies in the range of 2713.2 - 2927.7 A reduced in vacuum. The analysis was made on the basis of the similari-

ty of the bands of CCl and those of the isoelectronic

molecules CF, SiF, SiCl on the following assumptions: 1) The investigated bands belong to a $\sum -2\pi$ -transition; 2) the ground state of CCl is normal as in the case of CF, SiF and SiCl; 3) on the basis of the general process of the increasing doublet separation in the series SiF - SiCl - SiBr it may be expected that the doublet separation will in the case of CCl

be 1.5 - 2 times that of CF (77 cm⁻¹), i.e. approximately

130 cm⁻¹; 4) the distance between the atoms C and Cl in the diatomic molecule CCl was equated with the distance in CCl₄;

Card 1/2

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On the Spectrum of the CC1-Molecule

SOV/156-59-2-2/48

herefrom follows a rotation constant in the ground state of the CCl-molecule of approximately 0.6 cm 1; 5) following Venkatesvarlu the most intensive group of bands belongs to the sequence $\Lambda v = 0$. On the basis of the expressions given by R. Mulliken (Ref 4) for the terms of the upper and lower electron state the formula was derived for the P1-, P2-, Q_1 and Q_2 -branches of the $^2\Sigma$ - $^2\eta$ -transition. The measured distances between the individual bands were in good agreement with the calculated values. The formula for the Q-edges was set up. The doublet splitting amounted to A = 118 cm-1. The disnociation energy of the ground state of the CC1-molecule calculated by means of linear extrapolation proved to be approximately equal to 4.5 ev. There are 1 figure, 2 tables,

PRESENTED BY: Kafedra fizicheskoy khimii Moskovskogo gosudarstvennogo universiteta im. M. V. Lomonosova (Chair of Physical Chemistry, Moscow State University imeni M. V. Lomonosov)

SUBMITTED:

July 11, 1958

Card 2/2

24(7) 50V/156-59-2-3/48

AUTHORS: Kuzyakov, Yu. Ya., Tatevskiy, V. M.

TITLE: On the Spectrum of the Molecule CC1 (O spektre molekuly CC1)

PERIODICAL: Nauchnyye doklady vysshey shkoly. Khimiya i khimicheskaya

tekhnologiya, 1959, Nr 2, pp 237-239 (USSR)

ABSTRACT: The authors investigated the luminescence spectrum of an

electric discharge in carbon tetrachloride vapor. On this occasion they found the bands described by Barrow (Ref 1), however, in higher number (10 instead of 4) within the range 2337 - 2341 Å. The method of the experiments carried out by the authors is distinguished from Barrow's method by the fact that the authors used helium as carrier of the electric digcharge. The bands were found to belong to the molecule CCl as the oscillation constants (Table 1) calculated for this molecule according to the Deslaudres formula were in good agreement with the experimental values. The ionization potential of the CCl-molecule was determined to amount to 9.5 ev which agrees well with the ionization potentials of CO and Op

with respect to the order of magnitude. There are 2 tables

and 2 references, 1 of which is Soviet.

Card 1/2

On the Spectrum of the Molecule CC1+

SOV/156-59-2-3/48

PRESENTED BY: Kafedra fizicheskoy khimii Moskovskogo gosudarstvennogo

universiteta im. M. V. Lomonosova

(Chair of Physical Chemistry, Moscow State University

imeni M. V. Lomonosov)

SUBMITTED:

July 11, 1958

Card 2/2

APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

SOV/70-4-2-8/36 Akishin, P.A., Naumov, V.A. and Tatevskiy, V.M. AUTHORS:

An Electronographic Investigation of the Structure of TITLE:

Molecules of the Halides of Gallium and Yttrium

(Elektronograficheskoye issledovaniye stroyeniya molekut

galogenidov galliya i ittriya)

PERIODICAL: Kristallografiya, 1959, Vol 4, Nr 2, pp 194-200 (USSR)

Investigations were made, as in the previous paper. ABSTRACT:

using the sector-photometric method and the new electrono-

graph at the MGU. The results were analysed by successive approximations and the method of radial

discributions. Theoretical and experimental distribution

curves are reproduced. Ga2Cl6 and Ga2Br6 were dimeric

with Ga-Ga distances of 3.28 and 3.41 Å, respectively, and angles X_3 -Ga₂- X_4 of 112° + 3 and 110° + 3 and

 $Ga_1-X_5-Ga_2$ of $91^{\circ}\pm 3$ and $93^{\circ}\pm 3$. All other distances are

tabulated. The other compounds GaF3, GaI3, YF3, YCl3,

YBr, and YI, were plane triangular molecules with Me-X

Cardl/2

SOV/70-4-2-8/36 An Electronographic Investigation of the Structure of Moleculus of the Halides of Gallium and Yttrium

distances of 1.88, 2.44, 2.04, 2.47, 2.63, 2.80 ± 0.05 Å, respectively. These observations contradict some by Brode (Ref 3). Ga_2F_6 molecules were present (as shown by mass spectrometry) to an extent of (1% and Ga_2I_6 molecules were present to about 8% in GaI_3 at the m.p. There are 5 figures, 3 tables and 13 references, 3 of which are Soviet, 10 English.

ASSOCIATION: Moskovskiy gosudarstvennyy universitet imeni M.V. Lomonosova (Moscow State University imeni M.V. Lomonosov)

SUBMITTED: July 15, 1958

Card 2/2

APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

23(SOV/77-4-3-3/16

AUTHORS: Galashin, Ye.A. and Tatevskiy, V.M.

TITLE: On the Phenomena of Photographic Reversal

I. The Effect of Bromides, Desensitizers, Oxidisers and Reducing Agents on Fhotographic Reversal

PERIODICAL: Zhurnal nauchnoy i prikladnoy fotografii i kinematogra-

fii, 1959, Vol 4, Nr 3, pp 175-182 (USSR)

ABSTRACT: The authors carried out experiments to demonstrate

the effect of surplus KBr, various oxidizers, desensitizers and reducing agents on KBr transparencies, which after treatment with one of these agents were exposed to the radiation of mixed red and violet light.

Diagram Nr 1 shows the scheme of the illuminator. Starting from the assumption that solarization and the

Herschel effect are intimately related / reference 13_7, the authors, in order to establish, if possible, an experimentally proved parallelism between these pheno-

Card 1/7 mena, exposed some transparencies to white light prior

APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

On the Phenomena of Photographic Reversal. I. The Effect of Bromides, Desensitizers, Oxidizers and Reducing Agents on Photographic Reversal

to the common procedure. The effect of surplus bromides in the emulsion on photographic reversal and
the Herschel effect under radiation is illustrated
in graphs 2-4. They show that an increase of the
bromide concentration results in a loss of light
sensitivity in the emulsion, and an intensification
of the above-mentioned phenomena. In one case (graph
3) the authors used only violet light, but the curve
did not change its basic character. Solarization
did not depend on the admixture of long-wave radiation. The experiments carried out with oxidizers
(quinone) and desensitizers (pinacryptol) showed the
same effects as the bromide experiments. An increase
in the oxidizer and desensitizer concentration resulted in a diminution of the maximum optical density of the produced image and a gradual moving of the

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former stability. On the basis of the identity of cond reversal reestablished itself and increased the light) was first partially resolved, but soon a setransparency had been previously exposed to white effect of red and violet light the latent image (the curves. Graph 10 is of special interest. Ander the a general characteristic of all these blackening the shift of the maximum towards longer exposures is crease in the value of maximum optical density and dium nitrite on the Herschel phenomenon. An ingraphic reversal. Graph 10 shows the effect of soare illustrated in graphs 7-10. Graphs 7-9 re-present the effect of the reducing agents on photo-(hydroquinone, sodium sulfite and sodium nitrite) results of the experiments with reducing agents mexime towards shorter exposures (graphs 5-6). The

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On the Phenomena of Photographic Reversal. I. The Effect of Bromides, Desensitizers, Oxidizers and Reducing Agents on Photo-

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APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

On the Phenomena of Photographic Reversal. I. The Effect of Bromides, Desensitizers, Oxidizers and Reducing Agents on Photographic Reversal

the mechanisms of solarization and the Herschel effect, the authors assume that the observed regularities in photographic reversal can be explained by the coagulation theory / reference 7.7. The surplus bromides in the layer, and the presence of oxidizers and desensitizers reduce the quantum yield of the silver photolysis and hamper the development of new silver "germs". This circumstance creates favorable conditions for the processes of coagulation and recrystallization of the silver particles, which in this way lose their catalytic activity. The processes are characterized by resolution of the minute active nuclei, which yield to the growth of a small number of large but inactive particles. Metallic silver set free during photolysis does not create new nuclei, but deposits on already formed silver particles. As a result of the reduction

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On the Phenomena of Photographic Reversal. I. The Effect of Bromides, Desensitizers, Oxidizers and Reducing Agents on Photographic Reversal

in the number of active nuclei reversal takes place, manifesting itself as solarization, as well as Herschel effect. The increase in bromide and desensitizer concentration additionally hampers the development of new nuclei and creates still more favorable conditions for the coagulation and recrystallization of the silver "germs". The reversal occurs sooner and in lesser optical densities. In this way the diminution of the maximum optical density and the shift of the maximum of the blackening curve towards shorter exposures can be explained. The presence of reducing agents in the layer has the opposite effect. Due to the increased rate of photolysis of the silver bromide, favorable conditions are given for the development of many active centers. Coagulation is hampered. Solarization is not observed at all or only later.

Card 5/7

On the Phenomena of Photographic Reversal. I. The Effect of Bromides, Desensitizers, Oxidizers and Reducing Agents on Photographic Reversal

The maximum of optical density is increased and the maximum is shifted towards longer exposures. The Herschel effect can be noticed only in the first moment, when the elimination of the catalytic activity (a result of the coagulation of the numerous silver centers formed during the exposure) cannot be compensated for by the formation of new nuclei due to the effect of the long-wave radiation. The authors quote the Soviet scientists Kravets and his collaborators / reference 19 7, Ye.A. Kirillov / reference 9 7 and K.V.Chibisov / reference 8 7 in support of their opinion concerning the character of the observed phenomena. There are 9 graphs, 1 diagram and 33 references, 20 of which are German, 8 Soviet, 4 English and 1 French.

Card 6/7

APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

CIA-RDP86-00513R001755110018-2 "APPROVED FOR RELEASE: 07/16/2001

SOV/77-4-3-3/16

On the Phenomena of Photographic Reversal. I. The Effect of Bromides, Desensitizers, Oxidizers and Reducing Agents on Photographic Reversal

ASSOCIATION: Moskovskiy gosudarstvennyy universitet im. M.V. Lomonosova, khimicheskiy fakul'tet (Moscow State University imeni M.V.Lomonosov, Chemistry Department)

SUBMITTED: 2 April, 1957

Card 7/7

CIA-RDP86-00513R001755110018-2" APPROVED FOR RELEASE: 07/16/2001

SOV/51-6-1-8/30

SERVICE CONTROLLER PROGRAMMENT PROGRAMMENT

AUTHORS:

Chumayovskiy, N.A., Tatovskiy, V.M. and Yur'yov, Yu.K.

TITLE:

The Absorption and Raman Spectra of Selemophene and Its Methyl Howelegass (Spektry peglesheheniya i spektry kembinatsionnogo

rasseyaniya selencfena iyego metilgomologov)

PERIODICAL: Optika i Spektroskopiya, 1959, Vol 6, Nr 1, pp 45-50 (USSR)

ABSTRACT:

The paper reports studies of the increred and ultraviolet absorption spectra and Raman spectra of selenophone (I), 2-methylselenophone (II), 3-methylselenophene (III), 2,3-dimethylselenophene (IV), 2,4-dimethylselenophene (V), 3,4-dimethylselenophene (VI): The Raman spectra were obtained on a Steinheil spectrograph (linear dispersion 0.1 mm/A) The absorption spectra in the infrared were obtained using an IKS-11 spectrometer with LiF (3300-6000 cm-1), NaCl $(660-3000 \text{ cm}^{-1})$ and KBr $(400-600 \text{ cm}^{-1})$ prisms. An infrared spectrometer VIKSM-3 with a NaCl prism was also used. The ultraviolet absorption spectra were obtained in isocotane using a SF-4 spectrophotometer. Measurements of the infrared spectra, obtained using the VIKSM-3 spectrometer, and of the ultraviolet spectra were carried out at the Optics Laboratory of I.N.E.O.S. of the Academy of Sciences of the U.S.S.R. Selenophene and its homologues studied in the present work

Card 1/3

SOV/51-6-1-8/30

The Absorption and Raman Spectra of Selenophene and Its Kethyl Homologues

had the properties given in a table on p 50. Fig 1 gives the Raman spectra of substances I, III and V (curves a, 6 and 6 respectively). Fig 2 gives the infrared absorption spectra of all the six substances studied and Fig 3 gives the corresponding ultraviolet absorption spectra. In all the substances (with the exception of VI) characteristic intense absorption bands appeared in the infrared between 1209 and 1250 cm-1. The coincidence of certain frequencies and the general similarity of the infrared absorption spectra of thiophene and selenophene and its homologues can be taken as confirmation of the plane structure of selenophene, which belongs to the C2x type of symmetry. The Raman and the infrared absorption spectra of selenophene agree with the results reported by Gerding et al. (Ref 1). The spectra of selenophene homologues show characteristic frequencies due to vibrations of the substituents (table on pp 46-47). The ultraviolet absorption spectra of selenophene and its homologues obtained in the region 2200-2800 A did not differ greatly between each other (Fig 3). The ultraviolet spectra of thiophene and its homologues behave in a similar manner. The authors

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APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

SOV/51-6-1-8/30 The Absorption and Raman Spectra of Selenophene and Its Methyl Homologues

suggest that the infrared absorption and the Raman spectra of selenophene and its homologues may be used for identification of these compounds. There are 3 figures, 2 tables and 11 references, 5 of which are Soviet, 3 French, 2 English and 1 Japanese.

SUBMITTED: March 31, 1958

Card 3/3

SOV/51-7-3-4/21

AUTHORS:

Kesler, Kh., Pentin, Yu.A., Treshchova, Ye.G. and Tatevskiy, V.M.

TITLE:

Investigation of the Infrared Absorption Spectra of Hydrocarbons at Various Temperatures Both in the Liquid and Solid Phases.

PERIODICAL: Optika i spektroskopiya, 1959, Vol 7, Nr 3, pp 301-310 (USSR)

ABSTRACT:

The paper reports a study of the infrared absorption spectra of nine hydrocarbons at temperatures from room temperature (liquid phase) and at low temperatures (solid phase). The study was undertaken to find out the changes in the spectra which occur on solidification. The hydrocarbons studied were normal alkanes (n-heptane, n-octane), branched alkanes (3-methylheptane, 2,3-dimethylheptane, 2,4-dimethylpentane, 2,5-dimethylheptane, 2,4-dimethylpentane, 2,5-dimethylheptane, 2,3-dimethylhexane) and branched alkanes (2-methylheptane-2, 3,3-dimethylheptane-1). All these hydrocarbons were prepared and their properties determined in outside laboratories (acknowledgments are made to Prof. R. Ya. Levina and to A.V. Iogansen in this connection). Table 1 gives the degree of purity, the melting and crystallization points and the refractive index at 20°C (np0) of the nine hydrocarbons listed above. The infrared spectra were recorded in the region from 700 to 1800 cm-1 by means of a two-beam infrared spectrometer IKS-2 with a NaCl prism. The optical slit width was 7-10 cm-1.

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SOV/51-7-3-4/21 Investigation of the Infrared Absorption Spectra of Hydrocarbons at Various Temperatures Both in the Liquid and Solid Phases

For liquids, cells with effective layer thickness from 0.03 to 0.48 mm were used. At low temperatures a special cell was employed; its construction ensured the constancy of the layer thickness of a liquid or solid in it. Measurements were carried out at temperatures from +20°C In order to obtain good crystals and to avoid vitrification, to -172°C. the hydrocarbons were cooled slowly. Figs 1-3 show absorption spectra of the nine hydrocarbons at various temperatures. Tables 2-4 give the observed absorption maxima for the liquid and solid phases. The results obtained show that in the case of 2,4-dimethylpentane and 2,5-dimethylhexane only one (the most symmetrical) isomer exists in the crystal phase, but more than one rotational isomer is present in the liquid phase. The authors suggest that only those substances crystallize out which have one rotational isomer of sufficiently high symmetry necessary to form a correct molecular crystal lattice. There are 3 figures, 4 tables and 10 references, 3 of which are Soviet and 7 English.

SUELITTED: November 26, 1958

Card 2/2

APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

507/51-7-4-5/32

AUTHORS:

Kuzyakov, Ya.Ya. and Tatevskiy, V.Li.

TITLE:

Rotational Structure of the (1--1) and (0--1) Bands in the Spectrum

of the Carbon Monochloride Molecule

PERIODICAL:Optika i spektroskopiya, 1959, Vol 7, Mr 4, pp 467-471 (USSR)

ABSTRICT: The spectrum of carbon monochloride (CC1) contains a group of tands in the region 2713-2927 Å; they belong to transitious of the $^2\Sigma$ - $^2\Pi$ type. In the existing literature (Refs 1-3) only the vibrational structure of these bands is discussed. The present paper gives the first rotational analysis of two bands in the CCl spectrum and determination of the rotational constants, corresponding to the ground and excited states. The band spectrum of CCl was obtained by means of a diffraction spectrograph DFS-3 (2 A/mm dispersion in the first order). Only two band sequences $(\Delta v = 0, +1)$ are suitable for the rotational analysis. Fig 2 gives the microphotogram of the $2\Sigma-2\Pi_3/2$ component of the (1--1) band. Values of the line frequencies in various branches and the combination ratios are given in Tables 1 and 2 (all values are in The calculated rotational constants were found to be

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50V/51-7-4-5/32

Rotational Structure of the (1--1) and (0--1) Bands in the Spectrum of the Carbon Monochloride Molecule

 $B_1^{\rm H}=0.6465\pm0.0032$ cm⁻¹ for the $^2\Pi_3/2$ state, and $B_0^{\rm i}=0.6589\pm0.0038$, $B_1^{\rm i}=0.6551\pm0.0026$ cm⁻¹ for the $^2\Sigma$ state. There are 2 figures, 2 tables and 6 references, 1 of which is Soviet, 3 3nglish, 1 Indian and 1 translation from English into Russian.

SUBMITTED: February 9, 1959

Card 2/2

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"APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2

AKISHIN, P.A.; NAUMOV, V.A.; TATRVSKIY, V.M.

Electron diffraction investigation of the structure of molecules of vaporous gallium, yttrium, lanthanum, and neodymium halides. Vest. Mosk. un. Ser. mat., mekh., astron., fiz., khim. 14 no.1: 229-236 159. (MIRA 13:8)

Chare Physical Chemistry Morcon State University imeni M.U Lemence J, Min. Ness cd. dssk.

1. Kafedra fizicheskoy khimii Moskovskogo universiteta.

(Halides)

"APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2

5(4),24(5),5(3)

AUTHORS: Tatevskiy, V. M., Papulov, Yu. G. SOV/20-126-4-37/62

TITLE:

Quantum-mechanical Foundation of a Formula for the Energy of Alkane Formation (Kvantovomekhanicheskoye obosnovaniye for-

muly dlya energii obrazovaniya alkanov)

PERIODICAL:

Doklady Akademii nauk SSSR, 1959, Vol 126, Nr 4, pp 823-826

(USSR)

ABSTRACT:

For alkanes C_nH_{2n+2} , which only contain bonds CC and CH the general formula may be derived for the energy of formation

from the atoms as follows:

= $\sum_{CC} R_{CC} + \sum_{CH} R_{CH} + \sum_{CC,CC} R_{CC,CC} + \sum_{CC,CH} R_{CC,CH} +$

+ $\sum_{\text{CH,CH}}$ RCH,CH

(10). The signs CC and CH in the first two

sums denote the summing up of all bonds CC and CH, respectively. The signs CC, CC; CC, CH; CH, CH in the last three sums denote the summing up of all bond pairs CC and CC, CC and CH, CH and CH, respectively. The values R_{CC,CC} may be divided as follows:

Card 1/4

1) Quantities R'CC.CC, which refer to two CC-bonds possessing

Quantum-mechanical Foundation of a Formula for the SOV/20-126-4-37/62 Energy of Alkane Formation

a common C-atom; 2) Quantities $R_{CC,CC}^{"}$, which refer to two CC-bonds separated by a CC-bond. $R_{CC,CH}$ and $R_{CH,CH}$ may also be divided in an analogous way:

$$\sum_{\text{CC,CC}}^{R} {}_{\text{CC,CC}} {}_{\text{CC,CC}}^{R} {}_{\text{CC,CH}}^{R} {}_{\text{CC,CH}}^{R} {}_{\text{CC,CH}}^{R} {}_{\text{CC,CH}}^{R} {}_{\text{CC,CH}}^{R} {}_{\text{CC,CH}}^{R} {}_{\text{CH,CH}}^{R} {}_{\text{CH,CH}}^{$$

As a rule, the remaining terms of these equations (11) are smaller and may be neglected. On account of quantum-mechanical considerations the authors finally derived the following equation for the energy of formation $E_{C_nH_{2n+2}}$ of alkanes from

the atoms: $E_{C_nH_{2n+2}} = \sum_{i,j=1, i \in j}^{4} n_{ij}B_{ij}$ (16). The derivation of this equation is given and discussed in the present paper.

Card 2/4

Quantum-mechanical Foundation of a Formula for the SOV/20-126-4-37/62 Energy of Alkans Formation

stant and has the following value: $B_{ij} = R_{CC} + (\frac{4-i}{i} + \frac{4-j}{j})R_{CH} + (\frac{i-1}{2} + \frac{j-1}{2})R_{CC}, cc_{+}^{+}[(4-i) + (4-j)]R_{CC}, cH^{+} + [(\frac{3-j}{2})(\frac{4-j}{i}) + (\frac{3-j}{2})(\frac{4-j}{j})]R_{CH}^{i}, cH^{+} + \frac{1}{3} \sum_{i,j=1,i < j} n_{i,j}(i-1)(j-1).$ $\cdot \begin{bmatrix} R_{CC}^{t}, cc_{+} + 2R_{CC}^{g}, cc_{+}^{-1} + \frac{1}{3} \sum_{i,j=1,i < j} n_{i,j}(i-1)(4-j) + (j-1)(4-i) \end{bmatrix}.$ $\cdot \begin{bmatrix} R_{CC}^{t}, cH^{t} + 2R_{CH}^{g}, cH^{t} + \frac{1}{3} \sum_{i,j=1,i < j} n_{i,j}(4-i)(4-j) + (j-1)(4-i) \end{bmatrix}.$ $\cdot \begin{bmatrix} R_{CC}^{t}, cH^{t} + 2R_{CH}^{g}, cH^{t} + \frac{1}{3} \sum_{i,j=1,i < j} n_{i,j}(4-i)(4-j) \end{bmatrix} \begin{bmatrix} R_{CH}^{t}, cH^{t} + 2R_{CH}^{g}, cH^{t} + 2R_{CH}^{g}, cH^{t} \end{bmatrix}.$ $(17), \text{ where the quantities } f_{i,j}, \text{ with the exception of } f_{22}, f_{23}, \text{ and } f_{33}, \text{ are equal to zero.}$ $f_{22} = (\alpha_{22}^{-1/3}) \begin{bmatrix} R_{CC}^{t}, cc_{-}^{-R_{CC}^{g}}, cc_{-}^{-CR_{CC}^{g}}, cH^{t-2R_{CC}^{g}}, cH^{t-1}_{CH}, cH^{t-$

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Quantum-mechanical Foundation of a Formula for the SOV/20-126-4-37/62 Energy of Alkane Formation

 $f_{33} = (\alpha_{33}^{-1/3}) \left[R_{CC,CC}^{t} - R_{CC,CC}^{g} - 2R_{CC,CH}^{t} + 2R_{CC,CH}^{g} + R_{CH,CH}^{t} - R_{CH,CH}^{g} \right]$

Equation (16) is thus analogous to an equation already earlier derived by the authors (Ref 2), in which case they proceeded from the conceptions of the types and forms of bonds CC and CH; the equation is as follows:

 $E_{c_{n}H_{2n+2}} = \sum_{i,j=1,i \leqslant j}^{4} n_{ij}A_{ij}$ (19). A certain connection between the constants B_{ij} was determined. These problems will be dealt with separately. There are 2 references, 1 of which

ASSOCIATION: Moskovskiy gosudarstvennyy universitet im. M. V. Lomonosova (Moscow State University imeni M. V. Lomonosov)

PRESENTED: February 14, 1959, by V. I. Spitsyn, Academician

SUBMITTED: February 10, 1959

is Soviet.

Card 4/4

APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

TATEVSKIY, V.M.

PHASE I MOOK EXPLOITATION

807/4043

Tatevskiy, Vladimir Mikhaylqvich, Stanislav Semenovich Yarovoy, and Viktor Adol'fovich Benderskiy

Zakonomernosti i metody rescheta fiziko-khimicheskikh svoystv parafinovykh uglevodorodov(Regularities and Calculation Methods of Physicochemical Properties of Paraffin Hydrocarbons). Moscow, Gostoptekhizdat, 1960. 113 p. Errata slip inserted. 2,500 copies printed.

Exec. Ed.: L.A. L'vova; Tech. Ed.: I.G. Fedotova.

PURPOSE: This book is intended for synthesis chemists, engineers and scientific workers in the petroleum and petrochemicals industries, and aspirants and students working in the field of motor fuels and the chemistry of hydrocarbons.

COVERAGE: The book explains the methods for determining the basic physical and chemical properties of paraffin hydrocarbons (the alkanes), especially the high paraffins, which have been the subject of the most research in the last ten years. These methods make it possible to determine such properties of isomers of the higher hydrocarbons (C -C) as boiling point, density, molecular

Card 1/5

Regularities and Calculation Methods (Cont.)

SOV/4043

volume, molecular refraction, vapor pressure, heat evaporation, heat of combustion, and heat of formation. Tables of computed values of certain physical and chemical properties for the isomeric nonanes, decanes and undecanes are given. The researcher who has learned to use the analytical methods discussed in this book with respect to alkanes will be able to use similar-methods for determining the physical and chemical properties of other hydrocarbons alkanes, alkyds, alkadienes, alkylcyclanes, and alkylbenzols as well as other classes of organic compounds. Many properties of the C₅-C₁₀ alkanes and certain

properties of the C_{11} , have been determined by the authors. The author mentions

N. Hikitin, V.A. Ziborov, and A.A. Boldin. There are 54 references: 41 Soviet (including one translation), 1 German, and 12 English.

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, 4.	Heats of evaporation	•
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>//>	0-22-00

TATRYSKIY, V.M., prof., red.; L'VOVA, L.A., vedushchiy red.; MUKHINA, E.A., tekhn.red.

[Physicochemical properties of individual hydrocarbons; recommended values] Fiziko-khimicheskie svoistva individual'-nykh uglevodorodov; rekomenduemye znacheniia. Moskva, Gos. nauchno-tekhn.izd-vo neft. i gorno-toplivnoi lit-ry, 1960.

(MIRA 13:12)
412 p. (Hydrocarbons)

FROST, Andrey Vladimirovich, prof. [deceased]: Prinimali uchastiye:

BUSHMAKIN, I.N.; VVEDENSKIY, A.A.; GRYAZNOV, V.M.; DEMCET'YEVA,

M.I.: DIHTSES, A.I.; DOBROMRAVOV, R.K.; ZHARKOVA, V.R.; ZHERKO,

A.V.; IPAT'YEV, V.H.; KVYATKOVSKIY, D.A.; KOROBOV, V.V.; MOOR,

V.G.; NEMTSQV, M.S.; RAKOVSKIY, A.V.; REMIZ, Ye.K.; RUDKOVSKIY,

D.M.; RYSAKOV, M.V.; SEREBRYAKOVA, Ye.K.; STEPUKHOVICH, A.D.;

STRIGALEVA, N.V.; TATEVSKIY, V.M.; TILICHEYEV, M.D.; TRIFEL',

A.G.: PROST, O.I.; SHILTAYEVA, L.V.; SHCHEKIN, V.V., DOLGOPOLOV,

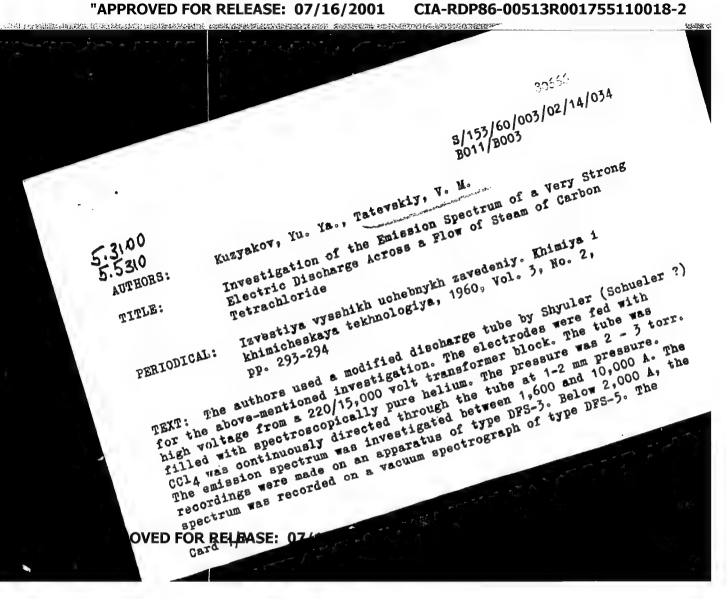
N.N., MOSTAVITED'; GERASIMOV, Ya.I., otv.red.; SMIRNOVA, I.V.; red.;

TOPCHIYEVA, K.V.; YASTREBOV, V.V., red.; KONDRASHKOVA, S.F., red.

izd-va; LAZAREVA, L.V., tekhn.red.

[Selected scientific works] Izbrannye nauchnye trudy. Moskva, #zd-vo Mosk.univ., 1960. 512 p. (MIRA 13:5)

 Chlen-korrespondent AN SSSR (for Gerasimov). (Chemistry, Physical and theoretical)



80663

Investigation of the Emission Spectrum of a Very Strong Electric Discharge Across a Flow of Steam of Carbon Tetrachloride

B/153/60/003/02/14/034 B011/B003

authors obtained the following results: 1. No spectra whatsoever were observed in the vacuum range and in the range of 7,000 - 10,000 A. 2. In the ultraviolet spectrum range the known groups of intense bands belonging to the CCl-molecule were obtained. Furthermore, R. E. Barrow's bands (2,300 A) were obtained (Ref. 1), which had been ascribed by him to CCl or CCl+. 3. In the range of 2,600 A a continuous background commences which reaches toward the direction of shorter wavelengths, i.e., to about 2,200 A. 4. In the visible and in the ultraviolet range numerous bands of the C2-molecule were obtained which belong to various transitions. 5. In the 4,050 A range a band emerges intensely belonging to the C3-molecule. The unclarified band system at 2,300 A is contrasted toward the direction of greater wavelengths and forms several sequences. The authors used the discharge of another type than that of Barrow and obtained ten bands instead of four, which belong to this system. The latter belongs to the CCl+-molecule. The authors calculated the molecular constants. The dissociation energy Do of the ground state of the CCl+-molecule is 51,700 cm-1. The

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Card 2/4

30534

Investigation of the Emission Spectrum of a Very Strong Electric Discharge Across a Flow of Steam of Carbon Tetrachloride

8/153/60/003/02/14/034 B011/B003

authors found that the oscillation analysis by P. Venkateswarlu (Ref. 2) contains contradictions and is unsatisfactory. A new oscillation analysis for the CC1-molecule which excludes the disadvantages mentioned in Ref. 2 (Refs. 3,4) was obtained by the authors from various analogies. They specify the oscillation constants obtained. Doublet cleavage A = 118 cm $^{-1}$. The authors refer to the table by Deslandres for the Q-edges of the CCl-molecule. The dissociation energy of the ground state of the molecule is ~4.5 ev on the strength of the linear extrapolation. Thus 2 band systems were detected in the ultraviolet range of the spectrum mentioned in the title. These systems are ascribed to the molecules CCl and CCl+. The nature of continuous emission between 2,200 - 2,550 A remained hitherto unclarified. In the discharge of the type used no chlorine bands could be observed. There are 1 table and 4 references, 2 of which are Soviet.

ASSOCIATION: Moskovskiy gosudarstvennyy universitet imeni M. V. Lononosova; Kafedra fizicheskoy khimii (Moscow State University imeni M. V. Lomonosov; Chair of Physical

Card 3/4

Chemistry)

Investigation of the Emission Spectrum of a Very Strong Electric Discharge Across a Flow of Steam of Carbon Tetrachloride

SUBMITTED: July 25, 1958

30666 8/153/60/003/02/14/034 B011/B003



Card 4/4

APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

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Card 1/2

s/051/60/009/002/007/013/XX E201/E491

Kuzyakov, Yu. Ya., Tatevskiy, V.M. and Tunitskiy, L.N. A Rotational Analysis of Boron Monoxide Bands Lying in AUTHORS:

PERIODICAL: Optika i spektroskopiya, 1960, Vol.9, No.2, pp.156-161 Chretien (Ref.1) studied the 1300 to 2100 Å spectra of discharges in BF3 with a vacuum spectrograph fitted with a grating of 1 m radius and 8.3 A/mm dispersion. Chretien found several bands which he ascribed to the BO molecule; the band edges, interpretations and relative intensities are given in Table 1. Later, Zelenskaya and Tunitskiy (Ref.2) reported a discharge Later, Letenskaya and Junitskiy (Ref. 2) reported a discharge spectrum of BF3 recorded in the vacuum ultraviolet region with a spectrograph Age -5 (DFS-5) with a resolving power of 120000 and a spectrograph Age -5 (DFS-5) with a resolving power of 120000 and a spectrograph Age -5 (DFS-5) with a resolving power of 120000 and a spectrograph Age -5 (DFS-5) with a resolving power of 120000 and the hands in the 1700 to 1000 A region to a hydrogen company of the bands in the 1700 to 1900 A region to a hydrogen compound of More detailed studies showed that Chretien's interpretation In the present paper the authors analyse the fine structure of the (0, 0) and (1, 0) bands (Chretien's nomenclature). A study of the isotopic shift in the BO bands, carried out in the authors' laboratory by A.A.Mal'tsev and V.M. Tatevskiy, showed that was correct. Chretien's interpretation should be corrected so that his (0, 0) and

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S/051/60/009/002/007/013/XX E201/E491

A Rotational Analysis of Boron Monoxide Bands Lying in the Vacuum Ultraviolet Region

(1, 0) bands become (1, 0) and (2, 0) respectively. These bands lie between 1300 and 2100 Å and are due to $2\pi \rightarrow x^2 \Sigma$ These two transitions of the BO molecule; interpretation of the (1, 0) band is given in Table 2 and of the (2, 0) band in Table 3. The rotational constants of the 21 state were also determined There are 4 tables and 7 references: 2 Soviet, 3 English, 1 Swiss and 1 translation from English into Russian.

November 23, 1959 SUBMITTED:

Card 2/2

CIA-RDP86-00513R001755110018-2"

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"APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2 5/051/60/009/006/004/018 Malitsev, A.A., Katayev, D.I., and Tatevskiy, V.M. An Investigation of the Electronic Spectral and the Isotopic Shift of Compounds of Boron and Oxygen. 24.6300 PERIODICAL: Optika i spektroskopiya, 1960, Vol.9, No.6, pp 713-720 AUTHORS: The electronic spectrum of BO has three band systems: The electronic spectrum of B0 has three band systems:
α-system in the 3100-8500 A region, β-system in the 2100-3600 A
region, β-system in the 3100-8500 A region, Chretien and Miescher unities
region, and γ-system discovered by Chretien and Miescher unities
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region, and γ-system discovered by Chretien and Miescher unities
Extending
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resolve
region, β-system in the B0 impurities
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region, β-system in the B0 impurities
region, β-system in the 2100-3600 Å
the isotopic shift of band edges in samples enriched with B10. TITLE: contradictions in interpretation of the BU Y-Dands by analysis the isotopic shift of band edges in samples enriched with Blo. the isotopic shift of band edges in samples enriched with Bio.

A discharge tube with hot hollow cathode was used to excite the spectra of Bo. A spectrograph in the 1600-2090 a region.

The spectra of Bo. A spectrograph in the 1600-2090 a region.

The wavelengths were found by comparing them with atomic lines of wavelengths were found by comparing them with atomic lines. the Y-bands of Bilo and blow in the low-20yu A region. It wavelengths were found by comparing them with atomic lines of boron, silicon and carbon. boron, silicon and carbon. Card 1/2

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An Investigation of the Electronic Spectra and the Isotopic Shift of Compounds of Boron and Oxygen. III. γ -Bands of the BO Molecule

the 1500-2090 Å region for Bllo (the upper spectrum) and Bl00 (the lower spectrum). Fig.2 shows the 0--3 and 1--4 bands of the γ-system of BO. Tables 2 and 3 list the wave-numbers (in cm-1) of the R2-branch band edges of Bllo (Table 2) and Bl00 (Table 3). The molecular potentials of BO are shown in Fig.3. It was found that the γ-system is due to the transition C2II--X2Σ. A more accurate vibrational analysis of the spectrum was derived from the results and the vibrational constants of Bllo and Bl00 were deduced (Table 4). A preliminary rotational analysis of five bands was carried out. Correlation between the electron states of BO and the atomic states of B and O was discussed. More accurate wavelengths of the atomic lines of boron and silicon (1360-2090 Å) were obtained; they are listed in Table 1. There are 3 figures, 3 tables and 31 references: 6 Soviet, 13 English, 4 German, 3 Dutch, 1 Swedish, 2 Swiss, 1 Indian, and 1 translation from English into Russian.

SUBMITTED: December 22, 1959

Card 2/2

KORYAZHKIN, V.A.; TATEYSKIY, V.M. Applicability of the ionic model to real diatomic molecules. Vest. Mosk un. Ser. 2: Khim. 15 no.4:21-22 JI-Ag *60. (MIRA 13:9) 1. Laboratoriya molekulyarnoy spektroskopii Moskovskogo universiteta. (Molecules)

"APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2

PAPULOV, Yu.G.; TATEVSKIY, V.M.

Energy of formation of a molecule as the sum of the energies of

paired atomic interactions. Vest. Mosk. un. Ser. 2: Khim. 15 no.5: 13-18 S-() 60.

1. Hoskovskiy gosudarstvennyy universitet, kafedra fizicheskoy khimii. (Molecules) (Heat of formation)

APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

"APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2

TATEVSKIY, V.M.; PAPULCY, Yu.G.

Relation between the energy of formation of a molecule from free atoms and its structure. Zhur. fiz. khim. 34 no.2:241-258 (MIRA 14:7) F '60.

1. Moskovskiy gosudarstvennyy universitet im. M.V.Lomonosova. (Molecules) (Atoms) (Chemical bonds)

APPROVED FOR RELEASE: 07/16/2001 CIA-RDP86-00513R001755110018-2"

AUTHORS:

s/076/60/034/03/001/038

TITLE:

Relationship Between the Formation Energy of a Molecule From Free Atoms and Its Structure. II. Energy of the Molecule as the Sum of Energies of the Interactions of Atoms in Pairs (Second Method)

Zhurnal fizicheskoy khimii, 1980, Vol 34, Nr 3, pp 489 - 504 (USSR)

TEXT: The fundamental concepts and equations for the relationship of the second PERIODICAL: method mentioned in the title and the fundamental approximate equation on the postulate of which the method is based are given. As a practical example, the formation energy of the alkane CnH2n+2 molecule is dealt with. The authors point out that equation (1) has no concrete meaning as long as it is not completed by an arbitrarily selected classification of the atom interaction in pairs. It may be seen from the symmetry of the ethane molecule (Fig 1) that all interactions in pairs $\eta_{\rm HH}$ are exactly equivalent with respects to energy, which does not hold for all interaction in pairs $\zeta_{
m HH}$. The same also applies to the interactions in pairs of the molecule of the optically active trans-isomer of n-butane (Fig 2). Bernstein's (D) classification, the classification (E), as well as the classification (G) developed by the authors are described, and the equations covering the Card 1/3

Relationship Between the Formation Energy of a S/076/60/034/03/001/038

Molecule From Free Atoms and Its Structure. II. Energy B115/B016

Molecule as the Sum of Energies of the Interactions of Atoms in Pairs (Second Method)

relationship mentioned in the title are derived. The interactions for various checkerboard configurations of the alkane molecule (Fig 3) as well as the classification of interactions in pairs (D) according to Bernstein are given in table 1. The stable checkerboard configurations of the atoms surrounding the C2 - C2, C2 - C3, and C3 - C3 bonds (the indices 2 and 3 designating in each case a secondary or tertiary carbon atom) are illustrated (Fig 4). The type of the C - C bonds and the position of six atoms surrounding the C - C bond in checkerboard equilibrium configurations are shown in figure 5. Equation (25) corresponds to Bernstein's classification and includes a superfluous constant. The mathematical equations used to calculate the coefficients of this equation by means of the number of different bonds in the alkanes are given in the paper. Classification (E) is defined with additional conditions, and the corresponding equations are derived. Classification (G) of the interactions of atoms in pairs developed by the authors, which is universally applied at present, is derived, and a corresponding equation for the formation energy of the alkane, which is equivalent to the equation given in the first method, is obtained. This classification

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Relationship Between the Formation Energy of a S/076/60/034/03/001/038 Molecule From Free Atoms and Its Structure. II. Energy B115/B016 of the Molecule as the Sum of Energies of the Interactions of Atoms in Pairs (Second Method)

is reproduced in table 2. From a comparison between the first and the second method it may be seen that, if a corresponding classification is used, each equation of the second method is mathematically fully equivalent to any equation of the first method with a corresponding classification of the bonds. A relationship can be established between the energy values of different bonds in the first calculation method and the energy values of different atom interactions in the second method. There are 5 figures, 2 tables, and 2 references, 1 of which is Soviet.

ASSOCIATION: Moskovskiy gosudarstvennyy universitet im. M. V. Lomonosova

(Moscow State University imeni M. V. Lomonosov)

SUBMITTED: February 16, 1959

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S/076/60/034/04/02/042 B010/B009

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AUTHORS:

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Tatevskiy, V. M., Papulov, Yu. G.

TITLE:

The Relation Between the Formation Energy of Molecules From

Free Atoms and Their Structure. III

PERIODICAL:

Zhurnal fizicheskoy khimii, 1960, Vol. 34, No. 4, pp. 708-715

TEXT: This is a discussion of the subject mentioned in the title. By way of introduction, it is pointed out that the attempt by several research workers to represent the formation energy of the molecules as the sum total of the bond energies and the energy of the interaction between the bonds runs counter to the generally accepted concepts of the structure of molecules. On the other hand, the equations derived in this way are not at all new. G. V. Bykov's (Ref. 5) observations on a relation between the bond energy and the "electron charges" of the bonds represent the laws established experimentally in a less acceptable way than do the equations given by the present authors in previous papers. The concept of "electron charges of bonds" given in Ref. 5 is a formal one and remains without foundation. A relation between the number of "effective charges" of the bonds and the bond energies is incompatible with the concepts of quantum

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The Relation Between the Formation Energy of Molecules From Free Atoms and Their Structure. III

S/076/60/034/04/02/042 B010/B009

mechanics. It is shown that the equations on the relation between the formation energy and the molecule structure of the alkanes derived according to the method described in a previous paper, are well able to represent the laws involved as well as some physicochemical properties (such as molar volume, molar refraction, heat of vaporization, etc.). These results are easily explained by and founded on the statements (Ref. 6) already made concerning the type and form of the C-C and C-H bonds. There are 26 references, 19 of which

ASSOCIATION: Moskovskiy gosudarstvennyy universitet im. M. V. Lomonosova (Moscow State University imeni M. V. Lomonosov)

SUBMITTED: February 16, 1959



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